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(54) Title: HUMANISED ANTIBODIES HAVING MODIFIED ALLOTYPIC DETERMINANTS

(57) Abstract

The invention relates to molecules which have an amino acid sequence derivable from part or all of the constant region of an immunoglobulin heavy chain. The constant regions are of a particular isotype and have one or more allotypic determinants. The amino acid sequence is substantially homologous to the amino acid sequence of the constant region. However, it has been altered so that it is without at least one of said allotypic determinants by making its sequence the site for an allotypic determinant identical to the amino acid sequence from the corresponding position in another equivalent immunoglobulin constant region of a different isotype. The invention provides synthetic immunoglobulins with reduced allotypic differences as compared to a given wild-type immunoglobulin.

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Humanised antibodies having modified allotypic determinants

The present invention relates to binding molecules.

In particular, it relates to recombinantly produced antibodies.

Owing to their high specificity for a given antigen, antibodies and particularly monoclonal antibodies (Kohler, G. and Milstein C., 1975 Nature 256:495) represented a significant technical break-through with important consequences scientifically, commercially and therapeutically.

Monoclonal antibodies are made by establishing an immortal cell line which is derived from a single immunoglobulin producing cell secreting one form of a biologically functional antibody molecule with a particular specificity.

Owing to their specificity, the therapeutic applications of monoclonal antibodies hold great promise for the treatment of a wide range of diseases (Clinical Applications of Monoclonal Antibodies, edited by E. S. Lennox. British Medical Bulletin 1984, publishers Churchill Livingstone). Antibodies are generally raised in animals, particularly rodents, and therefore the immunoglobulins produced bear characteristic features specific to that species. The repeated administration of these foreign rodent proteins for therapeutic purposes to

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human patients can 1 ad to harmful hypersensitivity reactions. In the main therefore, these rodent-derived monoclonal antibodies have limited therapeutic use. A further problem with these rodent derived antibodies, is that they are relatively ineffective at the depletion of cells in vivo, although the rat IgG2b antibody CAMPATH-1G is an exception to this rule.

which have characteristic features specific to human proteins. Unfortunately, immortal human antibody-producing cell lines are very difficult to establish and they give low yields of antibody (approximately 1 µg/ml). In contrast, equivalent rodent cell lines yield high amounts of antibody (approximately 100 µg/ml). Furthermore, where one wants to produce a human antibody with a particular specificity it is not practically or ethically feasible to immunise humans with an immunogen bearing the epitope of interest.

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In part, this problem has been overcome in recent years by using the techniques of recombinant DNA technology to 'humanise' non-human antibodies. Structurally, the simplest antibody (IgG) comprises four polypeptide chains, two heavy (H) chains and two light (L) chains inter-connected by disulphide bonds (see figure 1). The light chains are of two types, either kappa or lambda. Each f the H and L chains has a region

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f low sequence variability, the constant region (C) giving rise to allotypes and a region of high sequence variability, the variable region (V) giving rise to idiotypes. The antibody has a tail region (the Fc region) which comprises the C regions of the two H chains. The antibody also has two arms (the Fab region) each of which has a $V_{\rm L}$ and a $V_{\rm H}$ region associated with It is this pair of V regions $(V_{I_L}$ and $V_{H})$ each other. that differ from one antibody to another, and which together are responsible for recognising the antigen. In even more detail, each V region is made up from three complementarity determining regions (CDR) separated by four framework regions (FR). The CDRs are the most variable part of the variable regions, and they perform the critical antigen binding function. The CDR regions are derived from many potential germ line sequences via a complex process involving recombination, mutation and selection. It has been shown that the function of binding antigens can be performed by fragments of a whole Binding fragments are the Fv fragment which comprises the $V_{\rm L}$ and $V_{\rm H}$ of a single heavy chain variable domain (V_H) .

In creating "humanised" immunoglobulins, the Fc tail of a non-human antibody is exchanged for that of a human antibody. For a more complete humanisation, the FRs of the non-human antibody are exchanged for human FRs. This

process is carried out at the DNA lev l using recombinant techniques. However, these humanised immunoglobulins do not solve all the problems, because an immune response may still be mounted against the treatment antibody even when a patient is treated with a human antibody, as it may show certain sequence differences in the V (ie idiotypic differences) and C (ie allotypic differences) regions when compared with the patients own equivalent This is a particular problem where the antibodies. patient's immune system has already seen, and therefore been primed against, antibodies having these sequence differences (eg a patient may have received a prior blood transfusion which contained allotypically different immunoglobulins). A model system of injecting "mouseised human antibodies" into mice indicated that the allotype matching could critically affect the anti-idiotype response (Bruggemann M., Winter G., Waldmann H., Neuberger M.S., (1989) J. Exp. Med. 170, 2153-2157).

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The present applicants have realised that one way
around this problem is to eliminate the allotypic
variation from the constant region.

There are a range of different immunoglobulins IgG, IgM, IgA, IgD, IgE, known as isotypes, of which IgG is most commonly used therapeutically. It exists as isotypic sub-classes IgG1, IgG2, IgG3 and IgG4.

There are 24 recognis d allotypes of human

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immunoglobulin distributed between the different isotypes as follows:

IgG1 x 4

IgG2 x 1

IgG3 x 13

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IgA2 x 2

IgE x 1

Kappa x 3

The allotypes represent alternative amino acid substitutions found at discrete sites in the protein sequence. These different allotypic determinants are found in different combinations within given allelic forms of genes, but not all possible combinations which theoretically might exist are in practice observed.

For example, the four different allotypes of IgG1 can be seen (ie distinguished) by the immune system. These are Glm 1, 2, 3 and 17. Alternatively, combinations thereof, such as Glm (1, 17), can also be distinguished. The four different single allotypes are depicted in figure 2.

Antisera can be raised in other non-human species which can see the alternative isoallotypes provided that the antibody is purified away from the other human isotypes. Such isoallotypes for which such an antisera exists have been called non-allotypes and given the designation for example, nGlm(1) which is the isoallotype

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of Glm(1). Thus, although a human isoallotype should not be immunogenic in humans, it can still potentially be recognized in a different species.

Of the above mentioned different allotypes of IgG1, three common allelic forms of human IgG1 occur with different frequencies within different racial groups, namely Glm (3), Glm (1, 17), and Glm (1, 2, 17) based upon their reactivities with human antisera directed against the determinants Glm 1, 2, 3 and 17. At point in the future, it is likely that a patient with an existing anti-allotype response to one or more of these determinants will need treatment with a humanised The obvious solution and one which has been antibody. proposed in a letter to the Journal Nature (Mage, R.G., Nature (1988) 333, 807-808), is to make all the different allelic forms of an antibody and to allotype match each The present applicants have patient for therapy. realised that commercially this is not a good proposal because of increased production costs and the need to process several reagents in parallel through the Additionally, each patient regulatory requirements. would have to be tested for the response to different allotypes.

Thus, the present applicants propose eliminating the allotypes altogether from each therapeutic antibody. The sequence of the human allotype of IgG1 Glm (1, 2, 17) is

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shown aligned with sequences for the oth r human IgG, isotype sub-classes in figure 4 (a, b, c and d). It can be seen that the four isotypes are extremely homologous for the domains CH1, CH2 and CH3, and that the major isotypic differences are in the hinge region which varies in both, length and sequence between isotypes. The allotypic residues of IgG1 Glm (1, 2, 17) have been marked in figure 4. However, for the purposes of clarity the sequences around the allotypic sites Glm (1) (2) and (17) are shown below for each isotype.

Site (1)

	<u>355</u>	<u>356</u>	<u>357</u>	<u>358</u>	
	Arg	Asp or Glu	Glu	Leu or Met	IgG1
	Arg	Glu	Glu	Met	IgG2
15	Arg	Glu	Glu	Met	IgG3
	Gln	Glu	Glu	Met	IgG4

Thus, at site (1), IgG1 may exist as several allotypes depending on whether aspartic acid or glutamic acid at position 356, or leucine or methionine at position 358 are present.

Site 2

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	<u>430</u>	<u>431</u>	<u>432</u>	
	Glu	Gly or Ala	Leu	IgG1
	Glu	Ala	Leu	IgG2
25	Glu	Ala	Leu	IgG3
	Glu	Ala	Leu	IgG4

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Thus, at site (2), IgG1 may exist as ither of two allotypes depending on whether glycine or alanine is present at position 431.

Site (17)/(3)

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5 Sites (3) and (17) are alternative substitutions at the same site.

	<u>213</u>	214	215	
	Lys	Lys or Arg	Val	IgG1
	Lys	Thr	Val	IgG2
10	Lys	Arg	Val	IgG3
	Lys	Arg	Val	IgG4

Thus, at site (17)/(3), IgG1 may exist as either of two allotypes depending on whether lysine or arginine is present. The allotypes (17) and (3) cannot co-exist as they represent alternative substitutions at the same position.

The alternative alleles of Glm (1) and (2) do not provoke a human allotype response because of the homology of these alleles with the other IgG sub-classes in this region. These alleles are therefore called isoallotypes because they are only recognisable by xenoantisers (antisers from a different species) and only when the isotype is purified away from the other sub-classes.

Therefore, the present applicants propose the creation of a new IgG1 allele by site-directed mutagenesis of the gene, for exampl, an existing

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CAMPATH-1H monocl nal antib dy gene described below, so that the new allele consists entirely of isoallotypic determinants. The preparation of IgG1 mutants according to the teaching provided by the present applicants is shown schematically in figure 3.

For Glm (1) and Glm (2), the changes comprise simple substitution by the alternative isoallotypic residues. However, in the case of Glm (17) the conversion of lysine to arginine would in some cases merely change the allotype to an allotype that is recognised by certain individuals as a Glm (3) allotype despite the fact that this residue is homologous with IgG3 and IgG4. This apparent contradiction is thought to be because this arginine is seen in a tertiary epitope in the context of the other IgG1 specific residues in close proximity in the CH1 domain or hinge region. This indicates that in addition to changing lysine, other residues in CH1 or the hinge will need to be changed in order to create a new isoallotype.

Although the above and ensuing description is specifically directed to IgG1 and in particular, the CAMPATH-1H monoclonal antibody, the same approach can be used to create isoallotypes of the other human isotypes such as IgG2, IgG3 and kappa.

Thus, the present invention provides a first binding molecule derivable from a second binding molecule;

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which second binding molecule is an immun globulin, or a derivative, structural or functional analogue thereof, a member of a family of homologous molecules, and has one or more sites which are structurally distinctive from equivalent sites in the other family members;

wherein said first binding molecule is more closely homologous to the other family members than to said second binding molecule, at at least one of said one or more sites.

The first binding molecule may also be an immunoglobulin or a derivative, structural or functional analogue thereof. The one or more sites which are structurally distinctive from the equivalent sites in the other family members may be in the constant region giving rise to an allotypic difference. The first binding molecule may comprise entirely isoallotypic determinants.

The second binding molecule may be selected from the group consisting of IgG1, IgG2, IgG3, IgA2, IgE, kappa light chains or derivatives, structural or functional analogues thereof. Where the second binding molecule is IgG1, the allotypic differences may be present at one or more of sites (1) (2) (3) or (17) as described herein. Where the second binding molecule is IgG2, the allotypic difference may be present at site (23). Where the second binding molecule is IgG3, the allotypic differences may

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be pr sent at ne or more of th sites (11) (5) (13) (14) (10) (6) (24) (21) (15) (16) (26) or (27). Where the second binding molecule is IgA2, the allotypic differences may be present at one or more of the sites (1) and (2). Where the second binding molecule is kappa light chain, the allotypic differences may be present at one or more of the sites (1) (2) or (3). The sites referred to above are well documented in the literature (see e.g. Eur. J. Immunol. 1976.6:599-601. Review of the notation for the allotypic and related marks of human immunoglobulins).

The present invention also provides pharmaceutical preparations comprising a first binding molecule as defined above or described herein together with one or more excipients. The pharmaceutical preparation may comprise a cocktail of said first binding molecules.

Also provided by the present invention are methods for making a first binding molecule as defined above or described herein.

These methods comprise the steps of: a) identifying in said second binding molecule, one or more sites which are structurally distinctive from the equivalent sites in the other family members; b) making said first binding molecule whereby it is more closely homologous to the other family members than to said second binding molecule at at least one of said one or more sites.

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The first binding molecule may be made by providing a gene sequence encoding the second binding molecule and altering those parts of the gene sequence encoding said one or more sites. The gene sequence may be altered by site directed mutagenesis using oligonucleotide primers. The altered gene sequence may be incorporated into a cloning vector or expression vector. The expression vector may be used to transform a cell. The cell may be induced to express the altered gene sequence.

The present invention therefore provides cloning vectors and expression vectors incorporating the altered gene sequence. Also provided are cells transformed by expression vectors defined above. Also provided are cell cultures and products of cell cultures containing the first binding molecules. Also provided are recombinantly produced said first binding molecules.

Thus the present invention provides a molecule which comprises an amino acid sequence derivable from part or all of the constant region of an immunoglobulin heavy chain which constant regions are of a particular isotype and have one or more allotypic determinants

wherein said amino acid sequence is substantially homologous to the amino acid sequence of said constant region, but has been altered so that it is without at least one of said allotypic determinants, by making the amino acid residues at the site of an allotypic

d terminant identical to the amino acid residues from the corresponding position in another equivalent immunoglobulin constant region of a different isotype.

The molecule may comprise an amino acid sequence derivable from part or all of a human immunoglobulin constant region.

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The molecule may also comprise one or more polypeptides together with said amino acid sequence.

The polypeptide may comprise a functional biological domain. The domain may be such that it mediates any biological function. The functional biological domain may comprise a binding domain. The binding domain will have an ability to interact with another polypeptide. The interaction may be non-specific or specific.

The polypeptide, biological domain, binding domain and immunoglobulin-like binding domain may derive from the same source or a different source to the constant region.

The constant region may be from an immunoglobulin of
the isotype IgG. The isotype subclass may be IgG1 and
the molecule may no longer have one or more of the
allotypic determinants 1,2,3 and 17. The isotype
subclass may be IgG2 and the molecule may no longer have
the allotypic determinant 23. The isotype subclass may
be IgG3 and the molecule may no longer have one or more
of the allotypic determinants 11,5,13,14,10,6,24,21,15,

16,26 and 27.

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The constant region may be from an immunoglobulin of the isotype IgA2 and the molecule may no longer have either or both of the allotypic determinants 1 and 2.

5 The present invention also provides a pharmaceutical preparation which comprises a molecule as defined.

The present invention also provides a reagent which comprises a molecule as defined.

The present invention also provides a nucleotide 10 sequence encoding a molecule as defined.

The present invention also provides cloning and expression vectors comprising a nucleotide sequence as delivered above.

The present invention also provides host cells

comprising a cloning or expression vector as defined above.

The present invention also provides a method of preparing a molecule as defined above which comprises the steps of:

- 20 (a) identifying a constant region of an immunoglobulin heavy chain;
 - (b) comparing the identified constant region with constant regions from immunoglobulin heavy chains of the same isotype to locate allotypic determinants in the identified constant region;
 - (c) obtaining the coding sequence for the identified

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constant r gi n having all typic determinants;

- (d) altering the coding sequence so that it codess for a molecule without at least one of said allotypic determinants and by making the amino acid residues at the site for an allotypic determinant identical to the amino acid residues from the corresponding position in an equivalent immunoglobulin constant region of an isotype different to that of said identified constant region;
- (e) using said altered coding sequence in an expression system to produce a said molecule.

The present invention also provides a method of treating a patient which comprises administering a pharmaceutical preparation as defined above.

Of course, there are a number of different strategies which could be used in order to make the molecules with fewer allotypic determinants.

Genes encoding therapeutically useful antibodies are generally available in one of several different forms. They may be available as a cloned variable region DNA sequence with restriction sites at each end, suitable for recloning along with a chosen cloned constant region DNA sequence into a suitable expression vector. This is the strategy described herein for the constructs TF57-19, MTF121 and MTF123. Alternatively, they may be available as complete immunoglobulin DNA sequences including the V and C regions together, e.g. a cDNA clone of a complete

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humanised or human antibody.

Whatever the form in which the cloned immunoglobulion gene is obtained, the next step is to predict the amino acid sequence of the constant region from the DNA sequence. The DNA sequence can be obtained using a variety of strategies familiar to molecular biologists. The predicted amino acid sequence would then be checked for the amino acids known to vary as allotypes. Any isoallotypes present within the sequence can be left unaltered. Any allotypes present can be mutated.

The next step, is to decide what amino acid sequence to mutate the allotype to, in order to imitate an isoallotype. This is done by lining up the sequence with the corresponding region of the other immunoglobulin For all known allotypes, it has been found isotypes. that one or more of the other isotypes have invariant sequences for the homologous region. One of these sequences can then be chosen to form the basis for the changes to be made in the allotype in question. Having predicted the new amino acid sequence for the constant region, it is necessary to alter the existing DNA clone or to create a new DNA clone which will encode this Again there are several strategies available sequence. to molecular biologists in order to achieve this. In the case f the example CAMPATH-1H constructs described

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her in, the gamma-1 constant region was cloned in an M13TG131 single stranded phage vector. Mutagenic oligonucleotides were synthesised which were largely homologous to the single strand, but which contained base changes necessary to alter the codons for the critical amino acids. The mutagenesis was carried out using a commercial kit from Amersham International, High Wycombe, Bucks. Alternatively it would be possible to synthesise a complete artificial gene which encodes the predicted sequence.

Once mutated or newly synthesised, the gene is ready There are many different expression for expression. vectors available. Some of these are more suitable for expression in restricted cell types. Again it is within the standard technical expertise of one skilled in this field to choose and adapt expression vectors for this In the case of the CAMPATH-1H constructs described herein, modifications of the pSVgpt and pSVneo vectors have been used. These vectors have convenient cloning sites for the immunoglobulin variable and constant region, encoding DNA fragments adjacent to suitable promoter and enhancer sequences to allow expression in lymphoid cells. The vector allows the easy independent replacement of variable or constant region encoding DNA fragments. Thus, any suitable variable region can be subcloned into the vector, to give rise to

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a new specificity, or the variabl region can be kept and the constant region changed to give rise to a new isotype or allotype. Alternative vector systems are readily available.

Having removed allotypes from heavy chain constant regions by mutating them all to isoallotypes, it may still be desirable to consider the light chain effect in stimulating an immune response.

The most common kappa light chain allotype is Km(3)

in the general population. Therefore it may be sufficient to utilise this common kappa light chain allotype, as relatively few members of the population would see it as foreign.

Alternatively there are no lambda light chain allotypes. Therefore they could be used in combination with the de-allotyped molecules derivable from heavy chain constant regions.

Where one utilises the kappa light chain, the allotype Km(1,2) could first be mutated to the allotype Km(1). The light chain allotype Km(1) is often only weakly recognized in combination with certain heavy chain classes and subclasses, and so may not cause a major problem in therapeutic use.

In order that the present invention is more fully
understood embodiments will now be described in more
d tail, by way of example only, and not by way of

limitati n. Reference will be made (and has already been made in the text above) to the following figures in which:

figure 1 illustrates the structure of an IgG 5 antibody;

figure 2 shows the allotypes for the IgG1 antibody CAMPATH-1H;

figure 3 shows schematically the preparation of IgG1 mutants;

- figure 4 shows the IgG1 Glm (1,2,17) allotype sequence aligned to single allotypic examples of IgG2, 3 and 4 (none of these other subclasses have allotypic residues which cover the same residues as for the IgG1 allotypes);
- figure 5 shows the M13TG131 cloning vector containing the human gamma-1 constant region, showing cloning sites and modified polylinker;

figure 6 shows the original Hu4vH HuG1 pSVgpt expression vector and its modified version;

figure 7 shows the result of an ELISA assay testing different dilutions of the antibodies of mutants 1, 2 and wild type CAMPATH-1H for IgG1 specificity;

figure 8 shows the result of an autologous complement mediated lysis test using human peripheral blood lymphocytes; and

figure 9 illustrates an antibody-dependent cell-

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mediated cytotoxicity assay (ADCC) using CD3 activated interleukin-2 expanded human blastocytes cell effectors (E) and targets (T).

The starting antibody used for site-directed mutagenesis was CAMPATH-1H, a monoclonal antibody with a kappa light chain containing the human constant region sequence for IgG1 which carries the Glm (1, 17) allelic determinants. The whole IgG1 encoding region exists as approximately 2.3 kb HindIII-SphI restriction fragment cloned in an M13 vector. The M13TG131 cloning vector containing the human gamma-1 constant region showing cloning sites and modified polylinker is shown in figure 5.

The IgG1 encoding region is entered in the EMBL Sequence Database under the code number HS1GCC4. The accession number is AC J00228 (the printout from the database is provided herein as Appendix 1). sequence is for the Glm (1, 17) allotype. It covers 2009 bases from the 5' HindIII site (A)AGCTT including all of the coding region. It does not however, include some of the 3' non-coding region up to the SphI site. sequence provided by the EMBL Database is that of the upper strand of DNA. The CH1 domain starts at nucleotide 210 and ends at nucleotide 503. The mutagenic oligonucleotides MO1 and MO4 hybridise to nucleotides 486 The hinge region starts at nucleotide 892 and to 510.

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ends at nucleotide 936. The CH2 domain starts at nucleotide 1481 and ends at nucleotide 1803. The mutagenic oligonucleotide MO2 hybridises to nucleotides 1515 to 1543. The essential signal for the poly A tail is provided by nucleotides 1902 to 1908.

In M13TG131, the IgG1 coding region exists as a 2260 nucleotide fragment, of which the final 251 nucleotides are non-coding and therefore, inessential. Therefore, an embodiment of the invention could be carried out using the sequence information provided by the EMBL Sequence Database. It should be noted however, that the Sph1 restriction site referred to above, is present in the 3' end inessential non-coding region. Therefore, if the sequence data as provided by the EMBL database were being used, alternative restriction sites would have to be utilised.

Using site-directed mutagenesis, (carried out using protocols and reagents as supplied in kit form, Amersham code RPN. 1523, Amersham International Plc, Amersham, UK) the sequence corresponding to the Glm (1) allele was converted to the corresponding sequence found in the other sub-classes for IgG (Asp Glu Leu to Glu Glu Met at positions 356-358 in the CH3 domain).

The mutagenic oligonucleotides used were:

25 a) MO1 (to convert G1m (17) to G1m (3))
5' CTC TCA CCA ACT CTC TTG TCC ACC T 3';

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- b) MO2 (to convert Glm (1) t its isoallotype nGlm (1))
 5' GGT TCT TGG TCA TCT CCT CCC GGG ATG GG 3'; and
- c) MO4 (to eliminate Glm(3) by changing Lys to Thr in the CH1 region)
- 5' CTC TCA CCA ACA GTC TTG TCC ACC T 3'. The oligonucleotides were synthesised and then purified using an automated synthesizer and oligo purification columns supplied by Applied Biosystems (Applied Biosystems, 850 Lincoln Drive, Foster City, California, 94404 USA) following the manufacturers recommended Mutations were checked by Sanger Dideoxy protocols. sequencing (Sanger, F.S., Nicklen, S., and Coulson, A.R., (1977) Proc. Natl. Acad. Sci., USA, <u>74</u>, 5463) using As this newly formed standard protocols and kits. allotype sequence is found in all humans, there should be no immunological response to this alternative form of Glm Additionally and similarly, the lysine residue (1).responsible for the Glm (17) allotypic determinant was converted to an arginine residue corresponding to the Glm

The gene for this new constant region of mutant 1 carrying these three changes has been sequenced, incorporated into an expression vector containing the CAMPATH-1H V-region and expressed together with the CAMPATH-1H light chain which had been introduced by cotransfection.

allele (Lys 214-Arg; mutant 1).

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A further mutant has been made by replacing the critical arginine residue associated with Glm (3) of mutant 1 with a threonine residue, to produce a heavy chain which is the equivalent of IgG2 and which should fail to react with both anti-Glm (17) and anti-Glm (3) antisera (mutant 2).

Mutant 2 has also been sequenced, re-cloned in an expression vector containing the CAMPATH-1H light chain.

The supernatants of growing cultures containing

10 either of the two mutants were subsequently assayed for
the expression of a human IgG1 kappa product.

The mutations were introduced with the oligonucleotides listed above. The modified Hu4vHGlpSVgpt vector shown in figure 6 was used to simplify the subcloning of all the new mutants into the expression vector, owing to the possibility of use of two different "sticky ends" Bam HI and NotI. The expression vectors and $V_{\rm H}$ region sequences and expression, along with the light chains, in YO rat plasmacytoma cells are all as described in Riechmann L., Clark, M.R. Waldman H., Winter G. (1988) Nature 332, 323-327.

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From the positive cultures, the producers of the largest amount of the IgG1 product were selected to obtain purified antibody for biological assays to determine their allotypes and biological effector functions.

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Example 1

performed to verify that an IgG1 type antibody was produced by the mutants. This was tested with microtiter plates coated with anti-CAMPATH-idiotype antibody (YID 13.9). Wild type CAMPATH-IH antibody served as control. The bound antibody was detected with biotin-labelled anti-human kappa reagents or anti-human IgG reagent (monoclonals NH3/41 and NH3/130 respectively although other suitable reagents are commonly available) and subsequent development with streptavidin horseradish peroxidase. Figure 7 illustrates the results obtained for:

TF 57-19 ("wild type" CAMPATH-1H antibody, 0)

15 MTF 121 (mutant 1,Δ)

MTF 123 (mutant 2, 1)

and the wild type CAMPATH-1H (\P) in a known amount as standard. The concentrations had been estimated, and the starting dilutions adjusted to 50 µg/ml in PBS/10 mg/ml BSA. The starting dilution was used to prepare 8 two-fold dilutions.

The slope of the graph shows clearly that the CAMPATH-idiotype antibodies recognises mutants 1 and 2 to an extent equivalent to that of the wild type CAMPATH-1H, and that all three antibodies tested are present in similar concentrations as the standard.

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Example 2

The mutants' capability of autologous complement mediated lysis of human peripheral blood lymphocytes was tested.

Human peripheral blood mononuclear cells from a healthy donor were isolated from 60 ml defibrinated blood on a Lymphoprep* gradient (Nyeggard & Co., AS, Oslo, Norway). The cell pellet was washed in IMDM (Iscove's Modification of Dulbecco's Medium, Flow Laboratories, Scotland), and the cells were labelled with ⁵¹Cr. The starting dilution of antibodies used in the test was 50 µg/ml in PBS, 10 µg/ml BSA (dilution 1). Dilution 1 was further diluted 8 times two-fold to a final dilution of 1/128. Wild type antibody diluted in the same manner was used as a control.

The result is illustrated in figure 8. As can be seen, both antibody mutants show a very similar result in lysing the blood mononuclear cells as the wild type. The efficiency of the mutants is almost identical.

20 Example 3

Experiments were conducted to investigate whether or not, the mutant antibodies were capable of antibody-dependent cell-mediated cytotoxicity (ADCC) using CD3 activated interleukin-2 expanded human blastocytes as effectors (E) and targets (T). Cells were generated and used as both effectors and targets essentially as

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describ d in Riechmann L., Clark M.R., Waldmann H., Winter G., 1988, Nature 322, 323-327.

Preparation of Target Cells (T)

5 ml of blastocytes (3 x 10^6 cells) were labelled with 51_{CT} for 1 h. After 1 h the cells were washed and transferred in 6 equal aliquots in 100 µl IMDM 1% BSA, to 6 x 10 ml tubes containing 100 µl of the antibodies of mutants 1 and 2, and the control. The tubes were incubated for 1.5 h at room temperature. The cells were then washed with 10 ml IMDM 1% BSA and resuspended in 1.5 ml IMDM 1% BSA.

Preparation of Effector Cells (E)

Unlabelled blastocytes (2 x 10^6) were diluted 100:1 and 30:1 in IMDM 1% BSA medium. The ratios 100:1 and 30:1 refer to the final absolute ratios of effectors to 51 Cr labelled targets in the assay. Assays were performed in microtitre plates with a total volume of 200 µl per assay well. Thus 100 µl of targets at a concentration of 2 x 10^4 were put in each well ie 2 x 10^3 total cells. For E:T of 100:1, 100 µl of effectors at 2 x 10^6 were plated per well ie 2 x 10^5 . For E:T of 30:1 100 µl of effectors at 6 x 10^5 were put into each well ie

The efficiency percentage of specific ⁵¹Cr release was calculated as follows:

% specific 51Cr release =

(test release cpm - spontaneous (cpm) x 100

(total cpm - spontaneous cpm)

cpm = radioactive counts per minute as measured on a counter.

- The result is shown in figure 9. The figure shows that all of the antibodies tested released chromium. Wild type TF 57-19 and mutant 2 (MTF 123) released at about equal levels, whereas mutant 1 (MTF 121) shows a slightly higher release.
- These results clearly show that the mutants have biological activity comparable to the wild type CAMPATH1H antibody.

Example 4

The antibodies were tested in an assay specific for their Glm (3) allotypes reactivity using a monoclonal reagent from Oxoid (WHO/IVISS recognised agent, Study Code No HP 6027). These tests were performed in replicates of two.

Microtiter plates were coated with the anti-CAMPATH idiotype YID 13.9.4 antibody captive, and divided into three arrays of 4 x 4 wells. Into each of the three arrays, 4 x 5 fold dilutions of the antibody TF 57-19, MTF 121 and MTF 123 (50 µg/ml) in PBS 1% BSA and a control solution of PBS/BSA each were added.

25 After an incubation of 45 minutes at room temperature, the antibody solution was removed, and

- (i) to th first array was added a 1:500 dilution of biotin-labelled anti-Glm (3);
- (ii) to the second array was added a 1:100 dilution of biotin-labelled antibody (NH3/41) specific for the kappa light chain; and
- (iii) to the third array was added a 1:1000 dilution of biotin-labelled antibody (NH3/130) specific for human IgG1.

The microtiter plate was developed with streptavidin horseradish peroxidase.

The result is illustrated in Table 1. The numbers in the results represent the optical density (0.D) as measured in an ELISA plate reader multiplied by 100 ie 12 represents an 0.D of 0.12 and 70 an 0.D of 0.70.

The result clearly shows, that samples 1-3 all react with the antibodies specific for IgG1 (see also Example 1 above) and the kappa light chains. The control is negative. However, in the assay for Glm (3) specificity, only MTF 121 (mutant 1) shows reactivity, whereas the wild type TF 57-19, MTF 123 (mutant 2) and the PBS/BSA control did not show any response.

This result illustrates clearly that the elimination of sites recognised in the allotype response by site-directed mutagenis of these sites can overcome the problems of allotypic immuno-reactions. Although the examples refer to the mutagenesis of IgG1 only, it will

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be clear to the person skilled in the art that other immunoglobulin isotypes can be similarly modified.

Example 5

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The antibodies were tested in a conventional allotyping experiment using inhibition of red cell agglutination. The experiment was carried out using reagents supplied by the Central Laboratory of the Netherlands Red Cross, Blood Transfusion Service (PO Box 9190, 1006 AD Amsterdam, Netherlands).

Human blood group O Rhesus D red cells were washed and then aliquots separately labelled as described below with one of the following three relevant anti-RhD human sera having antibodies of known allotype.

- (1) anti-D Glm(az) = Glm(1,17)
- 15 (2) anti-D Glm(x) = Glm(2)
 - (3) anti-D Glm(f) = Glm(3)

Coating of Red Cells with Anti-Rh Antibodies

One volume of packed washed red blood cells were incubated with 4 volumes anti-Rh serum and 4 volumes (phosphate) buffered saline (PBS) at 37°C during 60 minutes. Every 15 minutes the cells were mixed by shaking.

After incubation the coated cells were washed four times with PBS and stored at 4°C in preservation fluid (although coated red blood cells can be stored at 4°C in PBS for one week).

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These coated red blood cells were then agglutinated with four antisera to the IgG1 allotypes as follows using the recommended dilution for each antiserum.

- (1) anti-Glm(z) = anti-Glm(17) 1 in 30 dilution
- (2) anti-Glm(a) = anti-Glm(1) 1 in 30 dilution
- (3) anti-Glm(x) = anti-Glm(2) 1 in 20 dilution
- (4) anti-Glm(f) = anti-Glm(3) 1 in 30 dilution

The wild-type CAMPATH-1H TF57-19 or the different CAMPATH-1H constructs (MTF 121, MTF 123) with the altered gamma-1 constant regions were then tested for their abilities to inhibit the agglutination of the red cells The inhibiting antibodies were by the above antisera. tried at concentrations of 0.5mg/ml, 0.25mg/ml and 0.125mg/ml in phosphate buffered saline containing 5% foetal bovine serum. Control sera containing IgG1 of allotype Glm(zax) or Glm(f) [Glm(1,2,17) or Glm(3)] were also included in the experiment and were used at dilutions of 1 in 10,20 and 40. Where it occurred the inhibition was most easily seen for the CAMPATH-1H antibodies at the 0.5mg/ml concentration and it was much weaker for 0.25mg/ml and no inhibition was seen at The control sera inhibited at all three 0.125mg/ml. The results for the highest dilutions tested. concentration are shown below.

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Allotype CAMPATH-1H constructs Control sera

WO 92/16562

		TF57-19	MTF121	MTF123	G1m(1,2,17)	G1m(3)
	G1m(1)	+	-	-	+	-
	G1m(2)	-	-	-	+	
	G1m(3)	-	+	-	-	+
5	Glm(17)	+		. •	+	-

The results are therefore consistent with the original wild type CAMPATH-1H antibody TF57-19 having allotype Glm(1,17). The new mutant MTF121 type as allotype Glm(3) whilst the mutant MTF123 fails to allotype for any of the IgG1 allotype markers Glm(1,2,3,17) i.e. it appears not to have an IgG1 allotype.

The skilled man will be able to use the binding molecules hereby provided to make pharmaceuticals according to standard techniques. Similarly the pharmaceuticals can be used in accordance with standard practices.

80	6	~		_	
0.08	28/23	26/23	31/35	15/16	
0.4	54/63	55/52	57/63	15/15	•
7	88/88	71/69	02/99	15/16	
10	. 61/66	71/68	67/73	15/17	
0.08	27/30	28/20	96/96	15/18	
0.4	45/47	48/41	50/55	18/18	
2	52/53	54/52	55/60	17/17	
10	52/59	59/53	56/58	15/16	
0.08	13/11	53/44	16/17	15/19	
0.4	13/12	64/65	16/16	15/18	
ત		69/69	15/17	15/16	
10	12/10	80/75	17/18	15/16	
Sample (dilution)	1) TF 57-19 (Wildtype)	2) MTF 121 (Mutant 1)	3) MTF 123 (Mutant 2)	4) PBS/BSA	
) 10 2 0.4 0.08 10 2 0.4 0.08 10 2 0.4	10 2 0.4 0.08 10 2 0.4 0.08 10 2 0.4 10 2 0.4 10 1 0.4 12/10 13/12 13/11 52/59 52/53 45/47 27/30 61/66 86/88 54/53	10 2 0.4 0.08 10 2 0.4 0.08 10 2 0.4 0.08 10 2 0.4 pe) 12/10 13/12 13/11 52/59 52/53 45/47 27/30 61/66 65/68 54/53 13/11 52/59 52/53 48/41 28/20 71/68 71/69 55/52	10 2 0.4 0.08 10 2 0.4 0.08 1 0 2 0.4 0.08 10 2 0.4 9 12/10 13/12 13/11 52/59 52/53 45/47 27/30 61/66 85/88 54/53 10/15 69/89 64/65 53/44 59/53 54/52 48/41 28/20 71/88 71/69 55/52 17/16 15/17 16/16 16/17 56/58 55/60 50/55 36/36 67/73 66/70 57/63	10 2 0.4 0.08 10 2 0.4 0.08 10 2 0.4 0.08 10 2 0.4 0.4 0.08 10 2 0.4 0.4 0.4 0.08 10 2 0.4 0.4 0.4 0.4 0.1 0.4 0.4 0.4 0.4 0.4 0.4 0.4 0.4 0.4 0.4

Table 1

SUBSTITUTE SHEET

APPENDIX 1 - Sheet (a)

```
HSIGCC4
          2009 bases
Human ig germline g-e-a region a: gamma-1 constant region
ID
     HSIGCC4
               standard; DNA; PRI; 2009 BP.
AC
     J00228;
     23-APR-1990 (reference update)
DT
     18-JUL-1985 (incorporated)
DT
DE
     Human ig germline g-e-a region a: gamma-1 constant
DE
     region
KW
     constant region; gamma-immunoglobulin; germ line;
     hinge exon; immunoglobulin; immunoglobulin heavy
KW
KW
     chain.
OS
     Homo sapiens (human)
OC
     Eukaryota; Metazoa; Chordata; Vertebrata; Tetrapoda;
OC
     Mammalia; Eutheria; Primates.
RN
     [1] (bases 1-2009)
     Ellison J.W., Berson B.J., Hood L.E.;
RA
RT
     "The nucleotide sequence of a human immunoglobulin
     c-gamma-1 gene";
RT
RL
     Nucleic Acids Res. 10:4071-4079(1982).
RN
     [2] (bases 469-1070, 1465-1821)
RA
     Takahashi N., Ueda S., Obata M., Nikaido T.,
     Nakai S., Honjo T.;
RA
RT
     "Structure of human immunoglobulin gamma genes:
RT
     Implications for evolution of a gene family";
RL
     Cell 29:671-679(1982).
CC
     [1] and [2] report that nucleotide divergence among
CC
     the four gamma genes is much greater in the hinge
CC
     regions than anywhere else. [2] also reports the
CC
     hinge regions of gamma-2, gamma-4, a gamma
CC
     pseudogene, and the 5' flanking, ch2, and ch3
CC
     domains of the gamma genes.
CC
CC
     this entry is part of a multigene region (region a)
CC
     containing the gamma-3, gamma-1, pseudo-epsilon, and
CC
     alpha-1 genes. see segment 1 for more comments.
     Key
                    Location/Qualifiers
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     CDS
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FT
                    /note="Ig gamma-1 heavy chain
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APPENDIX 1 - cont. Sheet (b)

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FT
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FT
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FT
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FT
                      /citation=([1],[2])
FT
                      /note="C in [1]; t in [2]"
\mathbf{FT}
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FT
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FT
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FT
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FT
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     CDS
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                      c-region hinge"
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     CDS
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                      c-region ch2 domain"
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                      /note="C in [1]; cc in [2]"
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FT
                      c-region ch3 domain"
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     conflict
FT
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\mathbf{FT}
                      /note="T in [1]; c in [2]"
FT
     Sequence 2009 BP; 418 A; 698 C; 566 G; 327 T; 0
S<sub>0</sub>
     Other;
SQ
```

APPENDIX 1 - cont. Sheet (c)

	•		
CAGGGAGGG GCTAAGGTGA CCCAGACACT GGACGCTGAA GCCCAGCTCT GTCCCACACC GGCCCATCGG TCTTCCCCCT	360 GCGGCGTGCA TGGTGACCGT AGCCCAGCAA TGTCTGCTGG	ccccccacr carecrcage cagecrager ccccraacc Tcccaagac cararccage crcrccacr ccrcager	960 GCCTCGCCCT GCCCCAGCCG GGGACCGTCA CCCTGAGGTC
S0 CAGGGAGGGG CCCAGACACT GCCCAGCTCT GGCCCATCGG	350 GCCCTGACCA CTCAGCAGCG GTGAATCACA GGAGGGAGGG	650 CCGCCCACT CATGCTCAGG CAGGCTAGGT GCCCTAACC TGCCAAGAGC CATATCCGGG CTCTCCACTC CCTCAGCTCG	950 GCCAGCCCAG CCAGGGACAG AACTCCTGGG TCTCCCGGAC
40 GGCTTTGGGG TGCCCATGAG CTGCGCCTGG CTCCACCAAG	GGTGTCGTG GAACTCAGGC AGTCCTCAGG ACTCTACTCC CCCAGACCTA CATCTGCAAC TTGGTGAGGG GCCAGCACAG CCTGGACGCA TCCCGGCTAT	640 GAGCCTCTGC TGGGCAGGCA GGCTCAGACC AAAGGCCAAA CTCCCAATCT	940 GCCCAGGTAA TAGCCTGCAT TCAGCACCTG ACCCTCATGA GACCCTGAGG
30 GCTGACCTT GCACACCCAA CCAGGGGCCT CTCTTGCAGC		630 TCTTCACCCG TCCCAGGCTC GCAGGTGCTG AGCCCACCCC	TGCCCACGT GCCCAGGTAA TGCCCTAGAG TAGCCTGCAT CATCTTCC TCAGCACCTG ACCCAAGGAC ACCCTCATGA GAGCCACGAA GACCCTGAGG
AGCTTTCTGG GGCAGGCCAG GGCAGGTGGC GCCAGCAGGT CCTCGCGGAC AGTTAAGAAC GCGGTCACAT GGCACCACCT GGCACCCTCC TCCAAGAGCA	320 GAACCGGTGA GCTGTCCTAC AGCTTGGGCA GACAAGAAAG	CAAGGCAGGC CCCGTCTGCC GAGAGGTCT TCTGGCTTTT CAGGCCCTGC ACACAAAGGG AGGACCCTGC CCCTGACCTA GACACCTTCT CTCCTCCCAG	920 AACTCACACA GCGGGACAGG CGTCCACCTC TCCCCCCAAA TGGTGGACGT
	310 CTACTTCCCC CACCTTCCCG GCCTCCAGC CACCAAGGTG	610 CAAGGCAGGC GAGAGGCTCT CAGGCCCTGC AGGACCCTGC	910 901 CTTGTGACAA 961 CCAGCTCAAG 1021 GGTGCTGACA 1081 GTCTTCCTCT
61 121 181 241	301 361 421 541 541	601 661 721 781 841	901 961 1021 1081

APPENDIX 1 - cont. Sheet (d)

	1210	1220	1230	1240	1250	1260
1201	1201 GACGGCGTGG AGGTGCATAA	AGGTGCATAA		TGCCAAGACA AAGCCGCGGG AGGAGCAGTA	AGGAGCAGTA	CAACAGCACG
1261	TACCGGGTGG	TCAGCGTCCT	CACCGTCCTG	CACCAGGACT	GGCTGAATGG	CAAGGAGTAC
1321	AAGTGCAAGG	TCTCCAACAA	AGCCCTCCCA	GCCCCCATCG AGAAAACCAT	AGAAAACCAT	CTCCAAAGCC
1381	Aaaggtggga	CCCGTGGGGT	GCGAGGGCCA	GCGAGGGCCA CATGGACAGA GGCCGGCTCG	GGCCGGCTCG	GCCCACCTC
1441		TGCCCTGAGA GTGACCGCTG		TACCAACCTC TGTCCTACAG GGCAGCCCCG AGAACCACAG	GGCAGCCCCG	AGAACCACAG
	1510	1520	1530	1540	1550	1.00
1501	1501 GTGTACACCC	TGCCCCCATC	CCGGGATGAG	CCGGGATGAG CTGACCAAGA ACCAGGTCAG	ACCAGGTCAG	CCTGAC
1561	CTGGTCAAAG	GCTTCTATCC	CAGCGACATC	CAGCGACATC GCCGTGGAGT GGGAGAGCAA	GGGAGAGCAA	TGGGCAGCCG
1621		GAGAACAACT ACAAGACCAC		GCCTCCCGTG CTGGACTCCG ACGGCTCCTT	ACGGCTCCTT	CTTCCTCTAC
1681	AGCAAGCTCA	CCGTGGACAA	CCGTGGACAA GAGCAGGTGG CAGCAGGGA ACGTCTTCTC	CAGCAGGGA	ACGICTICIC	ATGCTCCGTG
1741	Atgcatgagg	CTCTGCACAA	ATGCATGAGG CTCTGCACAA CCACTACACG CAGAAGAGCC TCTCCCTGTC	CAGAAGAGCC	TCTCCCTGTC	TCCGGGTAAA
	1810	1820	1830	1840		0
1801	TGAGTGCGAC	GGCCGGCAAG	1801 TGAGTGCGAC GGCCGGCAAG CCCCGCTCCC CGGGCTCTCG CGGTCGCACG AGGATGCTTG	CEGGCTCTCG	CGGTCGCACG	AGGATGCTTG
1861	GCACGTACCC	CCTGTACATA	GCACGTACCC CCTGTACATA CTÍCCCGGGC GCCCAGCATG GAAATAAAGC ACCCAGCGCT	GCCCAGCATG	GAAATAAAGC	ACCCAGGGGT
1921		CCTGCGAGAC	GCCCTGGGCC CCTGCGAGAC TGTGATGGTT CTTTCCACGG GTCAGGCCGA GTCTGAGGCC	CTTTCCACGG	GTCAGGCCGA	GTCTGAGGCC
1981	1981 TGAGTGGCAT GAGGGAGGCA GAGGGGT	สมออลออลอ	ン山でででしてませ			

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CLAIMS

1. A molecule which comprises an amino acid sequence derivable from part or all of the constant region of an immunoglobulin heavy chain which constant regions are of a particular isotype and have one or more allotypic determinants

wherein said amino acid sequence is substantially homologous to the amino acid sequence of said constant region, but has been altered so that it is without at least one of said allotypic determinants, by making the amino acid residues at the site of an allotypic determinant identical to the amino acid residues from the corresponding position in another equivalent immunoglobulin constant region of a different isotype.

2. A molecule according to claim 1 which comprises an amino acid sequence derivable from part or all of a human immunoglobulin constant region.

- 3. A molecule according to claim 1 or claim 2 which comprises one or more polypeptides together with said amino acid sequence.
- 25 4. A molecule according to claim 3 wherein the polypeptide comprises a functional biological domain.

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- 5. A molecule according to claim 4 wherein the functional biological domain comprises a binding domain.
- 5 6. A molecule according to claim 5 wherein the binding domain is an immunoglobulin-like binding domain.
- 7. A molecule according to claim 6 in which said immunoglobulin-like binding domain and said amino acid sequence are derivable from the same or different sources.
 - 8. A molecule according to any one of claims 1 to 7 wherein the constant region is from an immunoglobulin of the isotype IgG.
 - 9. A molecule according to claim 8 wherein the isotype subclass is IgG1 and the molecule no longer has one or more of the allotypic determinants 1,2,3 and 17.
 - 10. A molecule according to claim 8 wherein the isotype subclass is IgG2 and the molecule no longer has the allotypic determinant 23.
- 25 11. A molecule according to claim 8 wherein the isotype subclass is IgG3 and the molecule no longer has one or

more of the allotypic d terminants 11,5,13,14,10,6,24,21, 15,16,26 and 27.

- 12. A molecule according to any one of claims 1 to 7 wherein the constant region is from an immunoglobulin of the isotype IgA2 and the molecule no longer has either or both of the allotypic determinants 1 and 2.
- 13. A pharmaceutical preparation which comprises a molecule according to any one of claims 1 to 12.
 - 14. A reagent which comprises a molecule according to any one of claims 1 to 12.
- 15. A nucleotide sequence encoding a molecule according to any one of claims 1 to 12.
 - 16. A cloning or expression vector comprising a nucleotide sequence according to claim 15.

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- 17. A host cell comprising a cloning or expression vector according to claim 16.
- 18. A method of preparing a molecule according to any one of claims 1 to 12 which comprises the steps of:
 - (a) identifying a constant region of an immunoglobulin

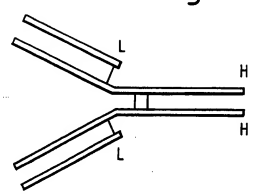
h avy chain;

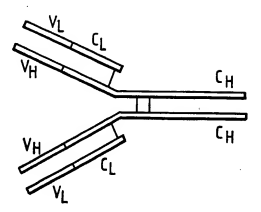
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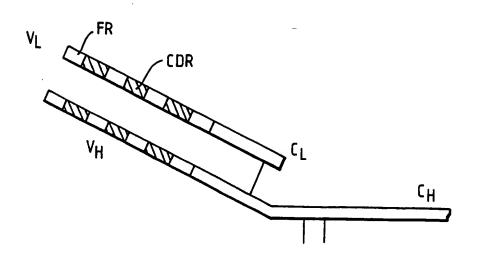
- (b) comparing the identified constant region with constant regions from immunoglobulin heavy chains of the same isotype to locate allotypic determinants in the identified constant region;
- (c) obtaining the coding sequence for the identified constant region having allotypic determinants;
- (d) altering the coding sequence so that it codes for a molecule without at least one of said allotypic determinants and by making the amino acid residues at the site for an allotypic determinant identical to the amino acid residues from the corresponding position in an equivalent immunoglobulin constant region of an isotype different to that of said identified constant region;
- (e) using said altered coding sequence in an expression system to produce a said molecule.
- 19. A method of treating a patient which comprises administering a pharmaceutical preparation according to claim 13.



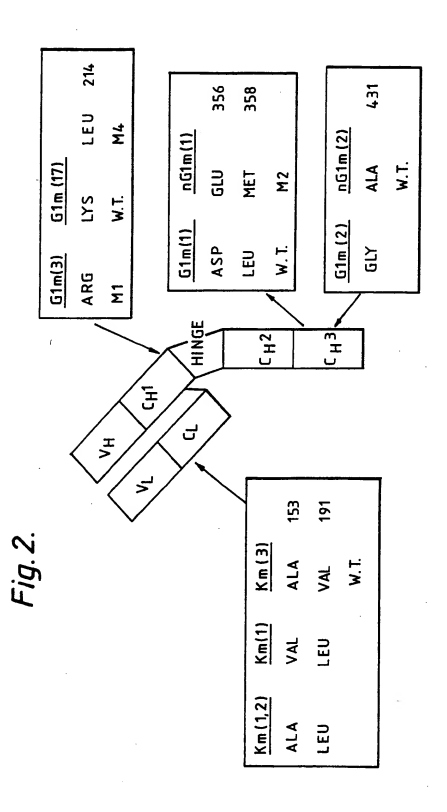
1∕11 Fig.1.







SUBSTITUTE SHEET



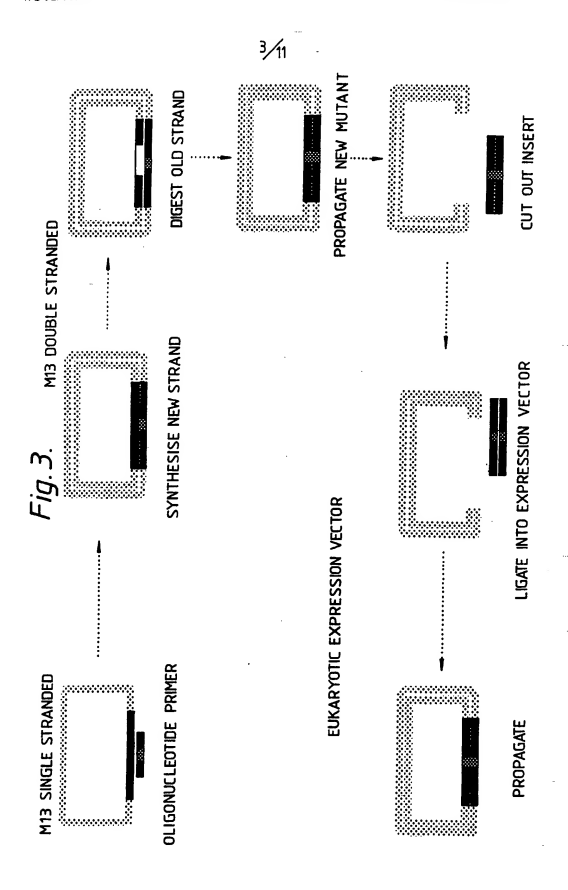


Fig.4a.

Human immunoglobulin sequences CH1 region

Ala	Ser	Thr	Lys	Gl	yPro	Sei	(Va	Pho	Pro	Lei	Mla	Pro	Ser	Se	Lys	Ser	Th	rSer	Gly	IgC1
=	-	-	-	-	-	-	-	-	•	-	-	-	_	-	Arg	_	-	-	Glx	IgC2
_	-	_	-	-	-	-	-	-	-	-	-	_	Cvs					-		
-	-	~	-	-	-	_	-	-	-	_	-	_								IgG4
													٠,٠		9					-804
Gly	Thr	Ala	Ala	Le	uGly	Cys	Leu	Va l	Lys	Asp	Туг	Phe	Pro	Cli	ıPro	Va 1	Th	rVal	Ser	IgGi
Ser	-	-	-	-	-	-	-	-	-	_	-	-	-	-	-	-	-	-	-	IgG2
-	-	-	-	-	-	-	-	-	-	_	-	-	-	-	_	_	_	-	_	IgG3
Ser	-	-	_	_	_	-	_	_	_	_	-	_	_	_	_	_	_	_	_	IgG4
													_		_	_				1604
Trp	As n	Ser	Gly	Al	aLeu	Thr	Ser	Gly	Val	His	Thr	Phe	Pro	Ala	Va 1	Leu	Gli	nSer	Ser	IgGi
-	-	-	-	•	•	-	-	-	-	-	•	-	•	-	-	-	-	-	-	IgG2
-	-	-	-	-	-	-	-	-	-	-	-	•	-	-	-	-	-	-	-	IgG3
-	-	•	-	-	-	•.	-	-	•	-	•	-	-	-	-	-	-	-	-	IgG4
Cly	Leu	Tyr	Ser	Lei	uSe r	Ser	Val	Va 1	Thr	Val	Pro	Ser	Ser	Ser	Leu	Cly	The	rGln	Thr	IgCi
-	-	-	-	-	•	•	-	-	-	-	•	-	- ,	Asn	Phe	•	-	-	-	IgC2
-	-	-	-	-	-	-	_	·_	-	-	-	-	-	-	-	-	_	-	-	IgC3
-	_	-	-	-	-	_	_	-	-	_	•	-	-	_	-	<u></u>	_	Lys	_	IgG4
																				-60
_		_					_	_							(C 1 m	(1)	")		
Tyr	lle	Cys.	Asn	Va.	lAsn	His	Lys	Pro	Ser	Asn	Thr	Lys	Va 1	Asp	Lys	Lys	Va.	l		IgC1
	Thr	•	-	-	Asp	•	•	•	-	-	•	-	-	-	- 3	Thr	-			IgG2
	Thr	•	-	-	Asp Asp	-	-	-	-	-	-	•	-	-	- ,	Arg	-			IgG3
- '	Thr	•	-	-	Asp	•	-	•	-	-	-	-	-	-	- ,	Arg	-			IgG4

Fig. 4b.

Human immunoglobulin sequences hinge region

GluProLys	SerCysAsp	LysThrH	lisTh	rCysProPro	IgGi
GlxArgLys				CysProPro	IgG2
				rCysProArgCysProGlu	IgC3
GluSerLysTyrGl				oCysProPro	IgC4
					IgG1
					IgG2
ProLysSerCysAs	ThrProPro	ProCysP	roAr	gCysProGluProLysSer	IgG3
		•			IgC4
				•	IgG1
					IgC2
CysAspThrProPro	ProCvsPro	ArgCvsP	roGl	uProLysSerCysAspThr	IgG3
•					IgC4
	CysPro				IgC1
	CysPro				IgG2
ProProProCysPro					IgG3
•	CysPro				IgG4

Fig.4c.

Human immunoglobulin sequences CH2 region

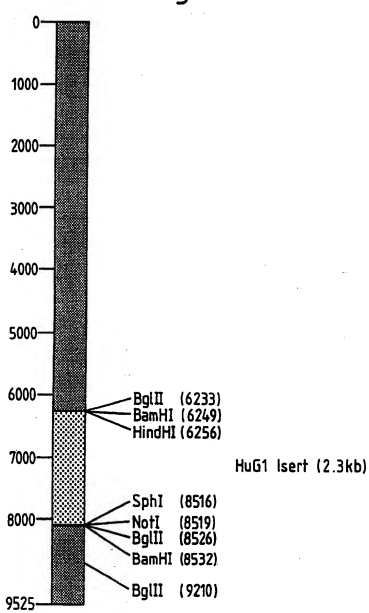
												Al				- BuLe		lyGl	y	IgC1
												-	-			1Al	4	-		IgG2
												-	-		•		•		- "	IgG3
												-	-	-	Pl	he -	•			IgC4
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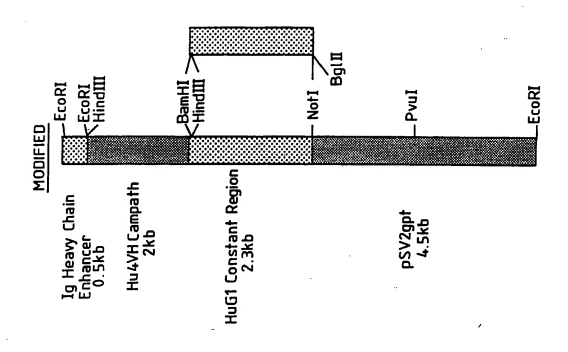
1∕11 Fig. 4d.

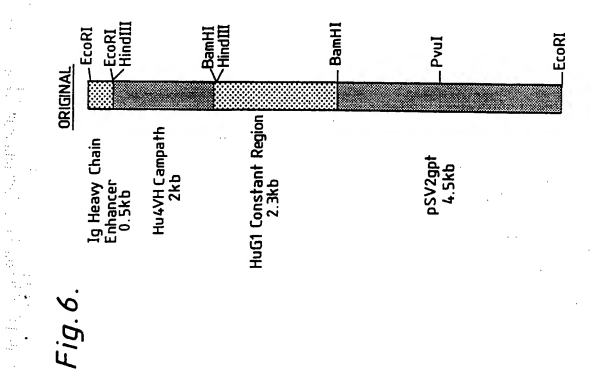
Human immunoglobulin sequences CH3 region

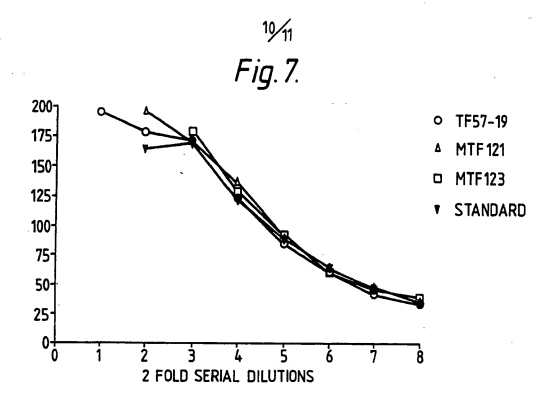
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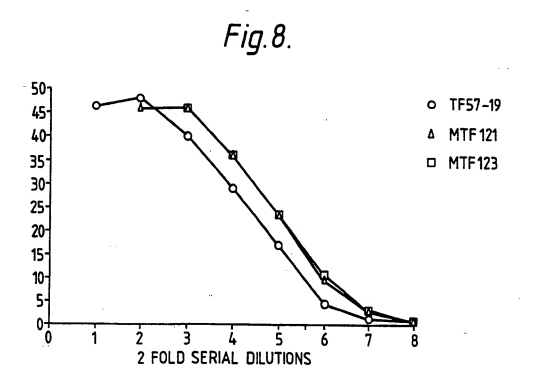
Fig. 5.



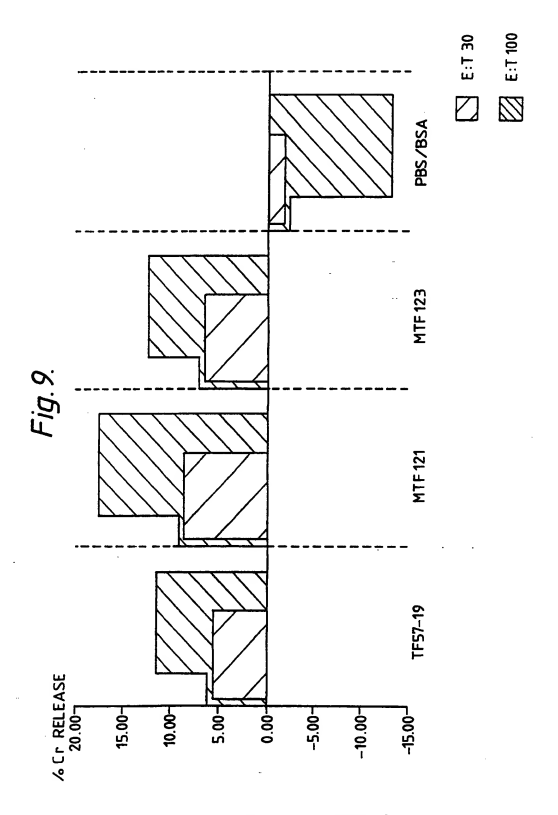








SUBSTITUTE SHEET



SUBSTITUTE SHEET

INTERNATIONAL SEARCH REPORT

International Application No.

PCT/GB 92/00445

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1. CLASSIFICAT	TON OF SUBJE	CT MATTER (if several classification	symbols apply, matche and	
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III. DOCUMEN	TS CONSIDERE	ED TO BE RELEVANT	6.4	Relevant to Claim No.13
Category °	Citation of D	ocument, 11 with Indication, where appro	printe, of the relevant passages	Junevalle to California
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"A" docume consider "E" earlier filling docume which is cristion other a "P" docume inter ti	red to be of parti- document but pul- are which may the se cited to establish or other special ent referring to a seens mt published prior the priority di- ATION	eneral state of the art which is not calar relevance blished on or after the international row doebts on priority claim(s) or the publication date of another reason (as specified) in oral disciosure, use, exhibition or or to the international filling date but	"I" inter document published after the in or priority date and not in conflict wichted to understand the principle or t invention "X" document of particular relevance; the cannot be considered novel or cannot involve an inventive step "Y" document of particular relevance; the cannot be considered to involve an indeciment is combined with one or meets, such combination being clovic in the art. "A" document member of the same patern. Date of Malling of this international	in the application but heavy underlying the a claimed invention to be considered to a claimed invention inventive step when the one other such documents to a person shilled at family
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International Application No

Page 2 PCT/GB 92/00445

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	W.E. PAUL, M.D.: "Fundamental Immunology", 1984,	1-19
١	Raven Press. New York. US; chapter 9: J.B.	į
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- International Ap atton No. PCT/ GB92/00445 FURTHER INF RMATI N CON ... AUED FROM THE SECOND SHEET V. X OBSERVATION WHERE CERTAIN CLAIMS WERE FOUND UNSEARCHABLE 1 This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons: 1. X Claim numbers Authority, namely: because they relate to subject matter not required to be searched by this Although claim 19 is directed to a method of the human body the search has been carried out and based on the alleged effects of the compositon. Claim numbers

because they relate to parts of the International application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically. because they are dependent claims and are not drafted in accordance with 3. Laim numbers the second and third sentences of PCT Rule 6.4(a). OBSERVATIONS WHERE UNITY OF INVENTION IS LACKING 2 This international Searching Authority found multiple inventions in this international application as follows: 1. As all required additional search fees were timely peld by the applicant, this international search report covers all searchable disims of the international application 2. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims of the international application for which fees were paid, specifically claims: No required additional search thes were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claim numbers: 4. As all searchable claims could be searched without effort justifying an additional fee, the international Searching Authority did not invite payment of any additional fee. Remark on Protest The additional search fees were accompanied by applicant's protest.

No protest accompanied the payment of additional search fees.

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ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

GB 9200445

SA 57491

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 23/06/92

The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

Patent document cited in search report	Publication date	Paten men	t family iber(s)	Publication date
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